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IMPORTANT SAFEGUARDS

READ ALL INSTRUCTIONS BEFORE OPERATING THIS DEVICE

NOTE, CAUTION AND WARNING STATEMENTS:

NOTE -Indicate some tips

CAUTION – Indicate correct operating or maintenance procedures in order to prevent damage to or destruction of the equipment or other property WARNING – Call attention to a potential danger that requires correct procedures or practices in order to prevent personal injury.



- 1. Always unplug this product immediately while it's not in use.
- 2. Do not disassemble the pump to avoid electrocution
- 3. Do not place or store product where it can fall or be pulled into a tub or sink.
- 4. Do not place in or drop into water or other liquid. Do not use while bathing.
- 5. Do not reach for a product that has fallen into water. Unplug immediately.

WARNING – To reduce the risk of burns, electrocution, fire or injury to persons

- 1. The operation of the system has to have the mattress connected to the PUMP, please do not power-off or unplug the PUMP in operation.
- 2. This product should never be left unattended when plugged in.
- 3. Close supervision in necessary when this product is used by, on, or near children or invalids.
- 4. Use this product only for its intended use as described in this manual. Do not use attachments not recommended by the manufacturer.
- Never operate this product if it has a damaged cord or plug, if it is not working properly, if it has been dropped or damaged, or dropped into water. Return the product to a service center or to the distributor for examination and repair.
- 6. Keep the cord away from heated surfaces.
- 7. Never block the air openings of this product or place it on a soft surface, such as a bed or couch, where the openings may be blocked. Keep the air opening free of lint, hair, and other similar particles.
- 8. Never drop or insert and object into any opening or hose.
- 9. Connect this product to a properly grounded outlet only. See Grounding Instruction.
- 10. Put the power cord or hose tube at the patient foot area to avoid wound on the patient's head.
- 11. To avoid electromagnetic interference, the patient environment should not have

- strong electro-magnetic or RF generated equipment near by
- 12. The PUMP will have minor heat generated in operation, please do not direct contact the surface continuously for more than 1 minute.
- 13. The product with ground pin (3 pin type) cannot be used in the home.
- 14. The EMC specification is compliant with the regulation requirement (please reference to the EMC information at the last page). For power cord with ground pin (3 pin type), the connection with properly grounded power outlet would get a better EMC suppressing effect. The system will work correctly also for the power cord connection with the power outlet without grounding.
- 15. When loss or failure of the supply mainstemporarily(In 20 minutes). It causespump stop, fail of power indication and alarm. But these are normal. The product can return to work state after supply mains is stable.

SYMBOLS	DESCRIPTION
I	POWER ON
0	POWER OFF
\triangle	ATTENTION
	DOUBLE ISOLATION
À	"BF" SYMBOL, INDICATE THIS PRODUCT IS ACCORDING TO THE
[X]	DEGREE OF PROTECTION AGAINST ELECTRIC SHOCK FOR TYPE BF
	EQUIPMENT
	CAUTION, READ THE INSTRUTION BEFORE USE
	AWAY FROM THE FLAME
IP21	WATER AND DUST PROTECTION CLASSIFICATION
TIA 250V	FUSE SPECIFICATION
	DISPOSAL OF ELECTRICAL & ELECTRONIC EQUIPMENT(WEEE): THIS
X	PRODUCT SHOULD BE HANDED OVER TO AN APPLICABLE
_	COLLECTION POINT FOR THE RECYCLING OF ELECTRICAL AND
	ELECTRONIC EQUIPMENT.
C. ASSIFE	UL CERTIFICATION LOGO (COMPLIACE WITH IEC60601-1)
_c ั(ปุ๊ _L) _{ับร}	With respect to electrical shock, fire and mechanical hazards only in
	accordance with STANDARD.
CB	CB CERTIFICATION LOGO



CE CERTIFICATION LOGO

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1 INTRODUCTION

This manual should be used for initial set up of the AKTC ELITE/SAVVY 8000 Series® Air Mattress System and for daily maintenance. Please keep the manual in handy area for reference.

2 INTENDED USE

This product is intended to help and reduce the incidence of pressure ulcers while optimizing patient comfort. It also provide following purposes:

- Individual home care setting and long-term care of whom suffering from pressure ulcer.
- Pain management as prescribed by physician.

⚠ NOTE: Equipment not suitable for use in the presence of a flammable anesthetic mixture with air or with oxygen or nitrous oxide.

3 PRODUCT DESCRIPTION

The ELITE/SAVVY 8000 Series are alternating mattress replacement system used in the prevention and treatment of pressure ulcers. By using the established principles of alternating therapy, the ELITE/SAVVY 8000 Series offer patients a comfortable and relaxing support surface which can both prevent skin breakdown and enhance healing.

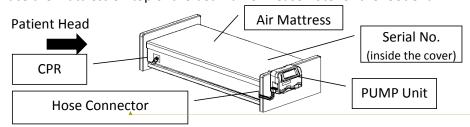
The CONTROL UNIT of the ELITE/SAVVY 8000 Series is a compact pump featuring an audible and visual low pressure, power failure and machine malfunction alarms, and a digital pressure adjustment function. The 19 cells mattress unit provides a unique design which keeps the lower layer of air cells constantly inflated while alternating and deflating the upper layer. The head section of cells remains static. The mattress has a heavy-duty nylon base sheet with a vapor permeable PU coated stretch cover.

In the event of cardiac arrest, rapid deflation is achieved by using the highly visible CPR facility.

4 PRODUCT INSTALLATION GUIDE

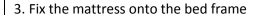
1. Unpacking the box to inspect for any damage, which may have occurred during shipment. If there are any damages, please contact your dealer immediately.

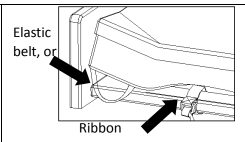




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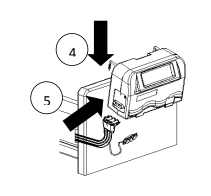
GDxxx-xxx/Rev.A

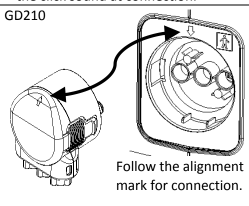


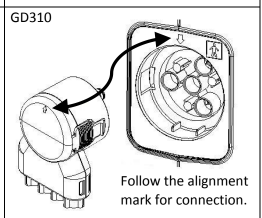


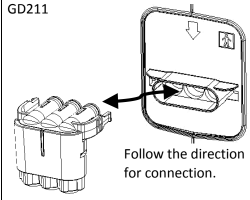
- 4. Hang the pump onto bed rail (food-end), the hangers will hold the bed rail tight automatically
- 5. Unplug the cover of the hose connector and connect the hose connector to the pump unit.

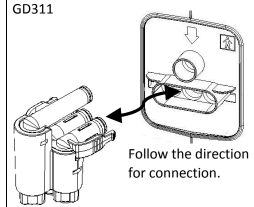
 Note: There's 4 combinations of hose connectors vs. the pumps'side panel as following (GD210, GD211, GD310, GD311), make sure you hear the click sound at connection.







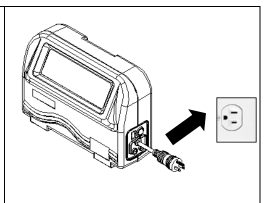




6. Plug the power cord into electrical outlet

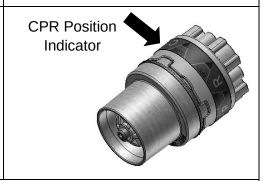
⚠ NOTE: Make sure the pump unit is suitable for the local power voltage

△ CAUTION: The pump can only be applied to the mattress recommended by the manufacturer. Do not use it for any other purpose (applied part: air

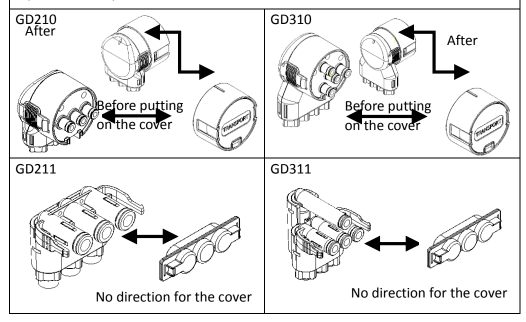


 Make sure the CPR is at CLOSE position before turning on the power.
 Switch the CPR to OPEN position to release the air at emergency or for packaging

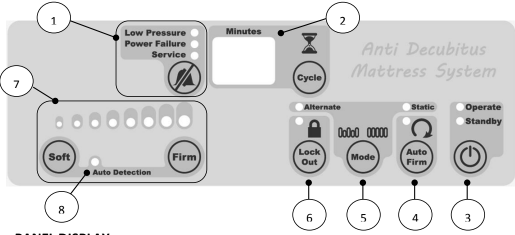
mattress)



8. Put on the hose connector cover at transportation, the mattress will retain pressure for up to 24hrs



5 PANEL DISPLAY AND THE OPERATION GUIDE



5.1 PANEL DISPLAY

- ① Alarm Mute and Alarm Indicator
 - Low Pressure Alarm Indicator
 - Power Failure Alarm Indicator
 - Service (Malfunction) Alarm Indicator
- ② Alternate Cycle Time or Warning codeDisplay
- ③ Operating or Standby
- (4) Auto-Firm
- (5) Function Mode Selection (Alternate & Static)
- (6) Panel Lock-out
- **7** Comfort Control
- **8** Auto Detection



5.1.1 ALARM MUTE

Press alarm mute button to temporary suspend the Low-Pressure/Power Failure/Static Overtime / Service alarms. Should the situation not resolved and the fault conditions continue, the alarm shall resume notifying the patient/caregiver.

5.1.2 Alternate CycleTime Display

Alternating Cycle Time can be selected from 10~30mins at 5mins interval by pressing the CYCLE button



5.1.3 Operate or Standby

Press this button to start operating or go into standby.

NOTE: The power switch on the side of pump must be turned on.At Power on the unit will resume the state before last power-off.



5.1.4 Inflate/Auto-Firm

The PUMP will go into the Inflate mode (LED lights flashing) every time theOPERATEmode is triggered. This insures the mattress to be able to reach its maximum operating pressure. Once the max pressure level is reached, the pump will automatically switch into the previous selected mode and comfort level. User can also use this function as full mattress inflation during patient sit-up or ingress/egress for better support.



5.1.5 Function Mode Switch

- ALTERNATE for the mattress to operate at alternating mode the air cell of the mattress will be proportionally deflated to reduce the surface pressure.
 The alternating cycle will continue at the selected cycle time until another mode is selected.
- STATIC This mode allows the mattress to maintain at the selected pressure.
 After 20 minutes, the STATIC OVER TIME alarm will be triggered for 10mins at every 15 seconds interval. Without further action, the PUMP will go into ALTERNATE mode automatically.



5.1.6 Panel Lock-Out

Should the panel remain untouched for 30 seconds or press the Lock-out button, the lock-out feature will lock the screen to prevent accident from changing the setting without notice. To unlock, press the Lock-out button for 3 seconds.



5.1.7 Comfort Level

Comfort level controls the air pressure output. When pressing the FIRM button,

GDxxx-xxx/Rev.A

the output pressure will increase and higher pressure output will support the heavier weight user, for decreasing air pressure, vice versa. Check to see if the suitable pressure is selected by sliding one hand between the air cells and the patient to feel patient's buttocks. Users should be able to feel the minimum contact. Always leave at least 1 inch space between user's buttock areas and air cells under to prevent bottoming out.



5.1.8 Auto Detection

When pressing the SOFT and FIRM button together, the pump will automatically detect the weight of the patient and set the appropriate pressureoutput for patient comfort.

5.2 OPERATION GUIDE

5.2.1 GENERAL OPERATION:

△ NOTE: The power switch is locatedon the side of pump

• Press to turn on the unit, all indicators on the control panel will light up accompanied with a beep for 2 seconds (You can also check the indicator for failure if any), and the indicator of STANDBY on the control panel will light up (In case the pump was turning off at OPERATE, it will go to OPERATE directly).

Ps: To test if the battery is working properly, press to turn off the power. Power failure alarm should be triggered. If not, please call customer service.

- Push on the OPERATE button , the system will start inflation and the
 "AUTO-FIRM" indicator will be flashing.
- Themattress shoud be fully inflated within 60 minutes, and automatically enter the last operating mode, otherwise the low pressure alarm will be triggered.
- According to the weight of the patient, adjust the pressure setting to the most suitable level without bottoming out. User can determine an appropriate pressure by adjusting the Comfort Level. Please consult with your physicianfor a proper setting.

Table 1 Weight and Common Level Melerence Table										
GDseries pump+8" mattress (iLAL/2-1Alternate)										
Comfort Control	Pump output			Pat	ient	Wei	ght (k	(G)		
(Auto-Detection)	Pressure(mmHg)	Pressure(mmHg) 20 40 60 80			100	120	140	160	180	
•••••	25	< 4	40							
•••••	30		20^	60						
•••••	35			40°	~80					
•••••	40				60~	100				
•••••	45					80~	120			
•••••	50						100^	⁻ 140		
••••••	55							120°	[~] 160	
0000000	60								140^	<mark>′180</mark>

Table 1 Weight and Comfort Level Reference Table

5.2.2 CPR

When CPR needs to be performed, quickly rotate the CPR valve to "OPEN" position, at the same time, disconnect the hose connector from the PUMP to speed up the air release.

5.2.3 AUDIBLE AND VISIBLE ALARM

(1) Power Failure – When electrical shortage occurred or power cord is unplug without turning off the pump, the "POWER FAILURE" indicator will light up along with buzzer. Check to ensure power cord is connected properly

 \triangle NOTE: When the PUMP is not been used for more than 3 months, it might need 6hours operating time or more for the Alarm to function properly.

- (2) Low Pressure –When an abnormal low pressure occurred in body section for4mins,the "Low Pressure" indicator will flash and beepevery 4 seconds. The Low Pressure alarm will continue until alarm mute button being pressed.Should the situation not be resolved and the fault conditions continue, the alarm will resume.
- (3) Static Overtime –When Static mode lasts for more than 20 minutes, "Static" indicator will flash and beep every 15 seconds. Press the MODE key to change to alternating mode can disable the alarm, or press the MUTE button to pause the alarm for 20 minutes. If not dealt, the system will automatically enter into alternating mode after 10 minutes.
- (4) Power Failure Overtime When the power is lost for more than 20mins, the "Power Failure" indicator will flash along with a beep sound at every 15 seconds, Press the MUTE button to cease the alarm, and check the patient for bed sore examination.

(5) Service(Malfunction)—When fault conditions occur, the "SERVICE" indicator will light up along with buzzer. Reference to Table 2 for Warning code and call the agent or distributor for service.

Table 2 WarningCode Reference Table

					TRETETICE TUDIC	
PRIORITY HIHG ↓ LOW	WARNINGCODE	INDICATOR LED	AUDIBLE OUTPUT MODE	CONDITION OF OUTPUT	WARNING DESCRIPTION	REMARKS
0	N/A	N/A	ONCE	Not in System Shutdown	Key Tone	Key Tone from Functional Button
1	S. d.	Power Failure	ONCE	POWER-OFF	System Shutdown	
2	8,8,	ALL LED	ONCE	OPERATE OR STANDBY	Power-On	All Indicators Light On
3	N/A	N/A	ONCE	OPERATE OR STANDBY	State/Mode Switching	
4	1. E.	AutoFirm	ONCE	OPERATE	Mattress Inflation Completion	Inflation Ended
5	R E	AutoFirm	ONCE	OPERATE	Auto-Firm Completion	Auto-Firm Ended
6	5, 8,	Static	ONCE	OPERATE	Static Completion	Static Ended
7	N/A	Power Failure	REPEAT (cycle4sec.)	POWER-OFF	Power Failure Alarm	No Display
8	I.F.	Low Pressure	REPEAT (cycle4sec.)	OPERATE OR STANDBY	Power-On Inflation FailureAlarm	
9	A.F.	Low Pressure	REPEAT (cycle4sec.)	OPERATE OR STANDBY	Auto-Firm Failure Alarm	
10	L.P.	Low Pressure	REPEAT (cycle4sec.)	OPERATE OR STANDBY	Low Pressure Overtime Alarm	
11	C.P.	Service	REPEAT (cycle4.5sec.)	OPERATE OR STANDBY	Constant Pressure Control Failure Alarm	
12	H.P.	Service	REPEAT (cycle4.5sec.)	OPERATE OR STANDBY	High Pressure Overtime Alarm	
13	L,E,	Service	REPEAT (cycle4.5sec.)	OPERATE OR STANDBY	Low Ambient Temperature Alarm	Environment Temperature Over Specification Limit
14	H]E,	Service	REPEAT (cycle4.5sec.)	OPERATE OR STANDBY	High Ambient Temperature Alarm	Environment Temperature Over Specification Limit
15	UI	Service	REPEAT (cycle4.5sec.)	OPERATE OR STANDBY	Air Valve 1 Positioning Failure Alarm	Air Valve 1 failure
16	U[2	Service	REPEAT (cycle4.5sec.)	OPERATE OR STANDBY	Air Valve 2 Positioning Failure Alarm	Air Valve 2 failure
17	P.E.	Power Failure	REPEAT (cycle15sec.)	OPERATE OR STANDBY	Power Failure Overtime Alarm	
18	L,b,	Service	REPEAT (cycle15sec.)	OPERATE OR STANDBY	Battery Low Alarm	Battery would need to be replaced
19	5,E,	Static	REPEAT (cycle15sec.)	OPERATE OR STANDBY	Static Overtime Alarm	
20	C,U,	NONE	NONE	FACTORY CALIBRATION MODE	Calibration Not Completed	
21	C,C,	NONE	NONE	FACTORY CALIBRATION MODE	Calibration Completed	

5.2.4 ALARM MUTE

When alarms were triggered, both the LED light and buzzer will sound off to warn the patient/caregiver. By pressing the button, it will temporary mute the buzzer so the caregiver may check for possible causes. Should the situation not resolved

andfault conditions continue, the alarm will resume. When in Power Failure situation, pressing alarm mute will cease the buzzer and turn off the "Power Failure" indicator.

6 CLEANING

By wiping the PUMP UNIT with a damp cloth pre-soaked with a mild detergent, and keep it away from dust. If other detergent is used, choose one that will have no chemical effects on the surface of the plastics case of the pump unit.

△ CAUTION: Do not immerse or soak pump unit.

By using a single use wipe, clean the MATTRESS COVER with a solution of neutral detergent and hand hot water. Rinse thoroughly with clean water and a damp single use wipe.

Disinfecting the cover

If the cover is heavily soiled or has been exposed to bodily fluids such as blood, it will require a more thorough cleaning procedure.

Wipe the cover using a single use wipe and a 0.1% Chlorine Solution (1,000ppm) and cold water. If required a 1% Chlorine Solution (10,000ppm) and cold water can be used. Rinse thoroughly with clean water and a damp single use wipe. Make sure the cover is completely dried before refitting to the mattress.

Frequent or prolonged exposure to higher concentration disinfectant solutions may prematurely age the fabric cover of mattresses. Surfaces must be protected during use and rinsed and thoroughly dried after application of a disinfectant.

Laundering

Before laundering mattress covers should be completely removed. Where required mattress covers can be laundered as follows:

Pre wash 60°C + 15 minutes

Main wash 60°C + 15 minutes

This should be followed by a cold rinse and extraction.

Drying

Mattress covers should be hung from a line or bar and drip dried in a clean indoor environment. Covers must be completely dried before refitting to the mattress.

Mattress covers can be tumble dried on a low heat setting for 90 minutes. Drying

temperature must not exceed 40°C . Exceeding the temperature can cause significant damage to the mattress cover.

 \triangle CAUTION: Do not use phenolic-based product for cleaning.

△ CAUTION: After cleaning, dry the mattress without direct exposure of sunlight.

7 STORAGE

- To quickly vacuum air out from mattress for storage, rotate the CPR valve to OPEN position and disconnect the hose connector to release the air.
- Lay the mattress out flat and upsides down.
- Roll from the head end towards the foot end
- Foot-end strap can then be stretched around the rolled mattress to prevent unrolling
- The power cord could be wrapped around the pump bumper or disconnected for storage.

8 MAINTENANCE

8.1 **8.1** General

- Check main power cord and plug if there are abrasions or excessive wears.
- Check mattress cover for signs of wear or damage. Ensure mattress cover and tubes are stubbed together correctly.
- Check the air hoses for any kink or break. For replacement, please contact your local dealers.

8.2 8.2 FUSE REPLACEMENT

- Disconnect the plug from mains power when a blown fuse is suspected.
- Remove the cover of the fuse holder by means of a small screwdriver.
- Insert a new fuse of the correct rating in, and replace the cover of the fuse holder back. The fuse rating should comply with the requested specification.

8.3 8.3 AIR FILTER REPLACEMENT

- Replace the air filter located at the back of the pump.
- The filter is reusable and can be washed gently with a mild detergent and water.
 Dry the filter before use.
- Check and replace air filter regularly if environment is dirty.

9 The Disposal of Air Mattress

When the air mattress is broken or no longer be useable, the mattress and the pump may be discard for recycle.

10 TROUBLESHOOTING

PROBLEM	SOLUTION
The mattress is not able to connect with the PUMP	 Check if the mattress model (model no. located inside the cover at the foot end)xxAAAxxx matched with the PUMP model xxBBB-xxx. The AAA should be the same as BBB. If not, please contact with the agent or distributor Check if the connector cover is removed and make sure the connector is not broken
The pump is showing no indications it is working	 Check if the plug is connected to mains Check if the main power switch is at ON position Check if there is any blown fuse
Power Failure Alarm Failure	The pump is in operation but the power failure alarm is not working at power down, please call customer service
The low pressure light is constantly flashing and the alarm is sounded	 Check if the CPR is at CLOSE position Check if the power was suddenly shut down Check if the connection between air tube to pump unit is tightly secured Check if all coupling connections along mattress are secured
Power Failure Alarm Failed	 If the PUMP is in operation but failed to trigger the Power Failure Alarm at Power Off, please contact the dealer or agent for further investigation
The pump is on but the mattress is not alternated	 Make sure the mattress inflationis completed Check the pump control panel the indicator of "ALTERNATE" is lighted on, if not, switch it to "ALTERNATE" Check if "Service" alarm indicator is on with buzzer, if yes, contact the dealer or agent for further investigation
The pump is operating noisily	 Make sure the pump is resting against a solid surface If the noise getting louder, contact the dealer or agent for further investigation
Patient is bottoming out (without alarm triggered)	Pressure setting might be inadequate for the patient, adjust comfort level to FIRM and wait for a few minutes for better comfort

If the above information does not solve the problem, please contact your localdealer or agent for further support.

11 TECHNICAL DATA

11.1 Product Specification

PUMP UNIT			AIR MATTRESS			
MODEL	GD Series Pump		MODEL	8" Mattress Series		
DIMENSION(cm)	33 (W) x 22 (D) x12 (H)	DIMENSION(cm)	89 (W) x 200 (L) x 21 (H)		
WEIGHT(kg)	3.5kg		WEIGHT(kg)	10Kg		
CYCLE TIME	10/15/20/25/30minut	tes	CELL MATERIAL			
STATIC TIME	30 minutes			Nylon TPU or TPU film		
AUTO FIRM TIME	20 minutes					
PUMP OUTPUT	> 8L (@120V or 230V)					
FLOW RANGE (Liter)	Note: The flow rate m	ay be				
	varied because of the					
	fluctuation of input vo	ltage				
PUMP OUTPUT			NO. OF AIR CELL			
PRESSURE RANGE	25 to 60 (±2)			19 CELLS		
(mmHg)						
POWER	AC120V 60Hz		COVER MATERIAL	2Way Stretched Polyester		
	or AC230 V 50Hz			with PU coated		
CURRENT	0.25 A _{MAX} (@132V~)		воттом	Nylon-TPU		
	or 0.12A _{MAX} (@253V~))	MATERIAL	Nylon-1PO		
FUSE RATING	T1AL 250VAC		MAX WEIGHT	180kg		
FREQUENCY	60Hz (120V)		MAX PRESSURE	103.5 mmHg		
	or 50 Hz (230V)			105.5 IIIIII g		
CLASSIFICATION	Class II					
	Type BF					
WARRANTY	2 years		WARRANTY	1 year		
SHELFLIFE	2 years		SHELFLIFE	1 year		
	ENVIRON	MENTA	L CONDITIONS			
ODEDATION ENVIRONM	AFNIT	5°C~40	0°C			
OPERATION ENVIRON	VIENI	15%RH ~ 93%RH(no condensation)				
		-25°&70°C				
STORAGE ENVIRONMENT		≤93%RH(no condensation)				
ENVIRONMENT PRESSURE		70 kPa-101.3kPa				
ENVIRONMENTHORIZONTAL LEVEL		≦3000m				
WATER AND DUST PRO	DTECTION	IP21				

11.2 EMC INFORMATION

Guidance and manufacturer's declaration-electromagnetic emissions

The <u>GD311-301</u> is intended for use in the electromagnetic environment specified below.

The customer or the user of the <u>GD311-301</u> should assure that it is used in such an environment.

Emission test	Compliance	Electromagnetic environment-guidance
RF emissions	Group 1	The GD311-301 uses RF energy only for its
CISPR 11		internal function. Therefore, its RF emissions
		are very low and are not likely to cause any
		interference in nearby electronic equipment.
RF emissions	Class B	The GD311-301 is suitable for use in all
CISPR 11		establishments, including domestic
Harmonic emissions	Class A	establishments and those directly connected
IEC 61000-3-2		to the public low-voltage power supply
Voltage fluctuations	Compliance	network that supplies buildings used for
/flicker emissions		domestic purposes.
IEC 61000-3-3		

Guidance and manufacturer's declaration-electromagnetic immunity

The <u>GD311-301</u> is intended for use in the electromagnetic environment specified below.

The customer or the user of the GD311-301 should assure that it is used in such an environment.

Immunity test	IEC 60601	Compliance level	Electromagnetic		
	test level		environment-guidance		
Electrostatic discharge(ESD)	± 6 kV contact	± 6 kV contact	Floors should be wood, concrete or		
IEC 61000-4-2	<u>+</u> 8 kV air	<u>+</u> 8 kV air	ceramic tile. If floors are covered		
			with synthetic material, the relative		
			humidity should be at least 30%		
Electrical fast transient/burst	± 2kV for power supply	± 2kV for power supply	Mains power quality should be that		
IEC 61000-4-4	lines	lines	of a typical commercial or hospital		
	± 1kV for input/output	Not applicable	environment.		
	lines				
Surge IEC 61000-4-5	± 1kV line(s) to line(s)	± 1kV differential mode	Mains power quality should be that		
	± 2kV line(s) to earth	Not applicable	of a typical commercial or hospital		
			environment.		
Voltage Dips, short	<5% UT(>95% dip in	<5% UT(>95% dip in	Mains power quality should be that		
interruptions and voltage	UT) for 0,5 cycle	UT) for 0,5 cycle	of a typical commercial or hospital		
variations on power supply	40% UT(60% dip in	40% UT(60% dip in	environment. If the user of the		
input lines IEC 61000-4-11	UT) for 5 cycles	UT) for 5 cycles	GD311-301		
	70% UT(30% dip in	70% UT(30% dip in	requires continued operation during		
	UT) for 25 cycles	UT) for 25 cycles	power mains interruptions, it is		
	<5% UT(>95% dip in	<5% UT(>95% dip in	recommended that the GD311-301		
	UT) for 5 s	UT) for 5 s	be powered from an uninterruptible		
			power supply or a battery.		
Power frequency(50/60 Hz)	3 A/m	3 A/m	The GD311-301 power frequency		
magnetic field IEC 61000-4-8			magnetic fields should be at levels		
			characteristic of a typical location		
			in a typical commercial or hospital		
			environment.		
NOTE UT is the a.c. mains voltage prior to application of the test level.					

Guidance and manufacturer's declaration-electromagnetic immunity

The GD311-301 is intended for use in the electromagnetic environment specified below.

The customer or the user of the GD311-301 should assure that is used in such and environment.

Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment-guidance
			Portable and mobile RF communications equipment
			should be used no closer to any part of the
			GD311-301 including cables, than the recommended
			separation distance calculated from the equation
			applicable to the frequency of the transmitter.
			Recommended separation distance:
			$d = 1,2 \sqrt{P}$
Conducted RF	3 Vrms	3 Vrms	d = 1,2 \(\sqrt{P} \) 80MHz to 800 MHz
IEC 61000-4-6	150 KHz to 80 MHz		$d = 2.3 \ \sqrt{P} \ 800 \text{MHz to } 2.5 \text{ GHz}$
Radiated RF	3 V/m	3 V/m	Where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in metres (m).
		3 V/III	
IEC 61000-4-3	80MHz to 2,5 GHz		Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, a should be less than the compliance level in each frequency range.
			Interference may occur in the vicinity of equipment marked with the following symbol:
			((·•))

NOTE1 At 80 MHz and 800 MHz, the higher frequency range applies.

NOTE2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

- a Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the GD311-301 is used exceeds the applicable RF compliance level above, the GD311-301 should be observed to verify normal operation. If abnormal performance is observed, additional measures my be necessary, such as re-orienting or relocating the GD311-301.
- Over the frequency range 150 kHz to 80 MHz, field strengths should be les than 3 V/m.

Recommended separation distance between portable and mobile RF communications equipment and the GD311-301

The <u>GD311-301</u> is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the <u>GD311-301</u> can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the <u>GD311-301</u> as recommended below, according to the maximum output power of the communications equipment.

Rated maximum output power of transmitter	Separation distance according to frequency of transmitter m				
W	150 kHz to 80 MHz	80 MHz to 800 MHz	800 MHz to 2,5 GHz		
	$d=1,2\sqrt{P}$	d=1,2√P	d =2,3√P		
0,01	0,12	0,12	0,23		
0,1	0,38	0,38	0,73		
1	1,2	1,2	2,3		
10	3,8	3,8	7,3		
100	12	12	23		

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in metres (m) can be estimated using the equation applicable to the frequency of the transmitter, where p is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

NOTE1 At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

NOTE2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

Guidance and manufacturer's declaration-electromagnetic emissions

The <u>GD311-401</u> is intended for use in the electromagnetic environment specified below.

The customer or the user of the <u>GD311-401</u> should assure that it is used in such an environment.

Emission test	Compliance	Electromagnetic environment-guidance
RF emissions	Group 1	The GD311-401 uses RF energy only for its
CISPR 11		internal function. Therefore, its RF emissions
		are very low and are not likely to cause any
		interference in nearby electronic equipment.
RF emissions	Class B	The GD311-401 is suitable for use in all
CISPR 11		establishments, including domestic
Harmonic emissions	Class A	establishments and those directly connected
IEC 61000-3-2		to the public low-voltage power supply
Voltage fluctuations	Compliance	network that supplies buildings used for
/flicker emissions		domestic purposes.
IEC 61000-3-3		

Guidance and manufacturer's declaration-electromagnetic immunity

The $\underline{GD311-401}$ is intended for use in the electromagnetic environment specified below.

The customer or the user of the GD311-401 should assure that it is used in such an environment.

Immunity test	IEC 60601	Compliance level	Electromagnetic
	test level		environment-guidance
Electrostatic discharge(ESD)	± 6 kV contact	± 6 kV contact	Floors should be wood, concrete or
IEC 61000-4-2	<u>+</u> 8 kV air	<u>+</u> 8 kV air	ceramic tile. If floors are covered
			with synthetic material, the relative
			humidity should be at least 30%
Electrical fast transient/burst	± 2kV for power supply	± 2kV for power supply	Mains power quality should be that
IEC 61000-4-4	lines	lines	of a typical commercial or hospital
	± 1kV for input/output	Not applicable	environment.
	lines		
Surge IEC 61000-4-5	± 1kV line(s) to line(s)	<u>+</u> 1kV differential mode	Mains power quality should be that
	± 2kV line(s) to earth	Not applicable	of a typical commercial or hospital
			environment.
Voltage Dips, short	<5% UT(>95% dip in	<5% UT(>95% dip in	Mains power quality should be that
interruptions and voltage	UT) for 0,5 cycle	UT) for 0,5 cycle	of a typical commercial or hospital
variations on power supply	40% UT(60% dip in	40% UT(60% dip in	environment. If the user of the
input lines IEC 61000-4-11	UT) for 5 cycles	UT) for 5 cycles	<u>GD311-401</u>
	70% UT(30% dip in	70% UT(30% dip in	requires continued operation during
	UT) for 25 cycles	UT) for 25 cycles	power mains interruptions, it is
	<5% UT(>95% dip in	<5% UT(>95% dip in	recommended that the GD311-401
	UT) for 5 s	UT) for 5 s	be powered from an uninterruptible
			power supply or a battery.
Power frequency(50/60 Hz)	3 A/m	3 A/m	The GD311-401 power frequency
magnetic field IEC 61000-4-8			magnetic fields should be at levels
			characteristic of a typical location
			in a typical commercial or hospital
			environment.
NOTE UT is the a.c. main	s voltage prior to application	n of the test level.	

Guidance and manufacturer's declaration-electromagnetic immunity

The GD311-401 is intended for use in the electromagnetic environment specified below.

The customer or the user of the GD311-401 should assure that is used in such and environment.

Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment-guidance
			Portable and mobile RF communications equipment
			should be used no closer to any part of the
			GD311-401 including cables, than the recommended
			separation distance calculated from the equation
			applicable to the frequency of the transmitter.
			Posterior de la constant de la const
			Recommended separation distance:
			$d = 1,2 \sqrt{P}$
Conducted RF	3 Vrms	3 Vrms	$d = 1.2 \ \sqrt{P} \ 80 \text{MHz}$ to 800 MHz
IEC 61000-4-6	150 KHz to 80 MHz		$d = 2.3 \sqrt{P}$ 800MHz to 2.5 GHz
Radiated RF	3 V/m	3 V/m	Where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in metres (m).
IEC 61000-4-3	80MHz to 2,5 GHz		Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, a should be less than the compliance level in each frequency range.
			Interference may occur in the vicinity of equipment marked with the following symbol:
			(((•)))

NOTE1 At 80 MHz and 800 MHz, the higher frequency range applies.

NOTE2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

- a Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the GD311-401 is used exceeds the applicable RF compliance level above, the GD311-401 should be observed to verify normal operation. If abnormal performance is observed, additional measures my be necessary, such as re-orienting or relocating the GD311-401.
- Over the frequency range 150 kHz to 80 MHz, field strengths should be les than 3 V/m.

Recommended separation distance between portable and mobile RF communications equipment and the <u>GD311-401</u>

The <u>GD311-401</u> is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the <u>GD311-401</u> can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the <u>GD311-401</u> as recommended below, according to the maximum output power of the communications equipment.

Rated maximum output power of transmitter	Separation distance according to frequency of transmitter m			
W	150 kHz to 80 MHz	80 MHz to 800 MHz	800 MHz to 2,5 GHz	
	$d=1,2\sqrt{P}$	d =1,2√P	d =2,3√P	
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10	3,8	3,8	7,3	
100	12	12	23	

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in metres (m) can be estimated using the equation applicable to the frequency of the transmitter, where p is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

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